The Claims:

This listing of claims will replace all prior versions, and listings, of claims in the application.

Listing of claims:

1. (Currently amended) A method of manufacturing an a waterinsoluble azole antifungal active agent-oral dosage form, said method comprising the steps of:

providing a single phase working solution comprising an a water-insoluble azole antifungal active agent, water, a water-soluble polymer and a solvent, said solvent selected from the group consisting of alcohol, acetone, and mixtures thereof; providing core particles formed from a pharmaceutically acceptable material; combining said working solution with said particles to produce—an a water-insoluble azole antifungal active agent-coated particles; drying said—an water-insoluble azole antifungal active agent -coated particles; and forming said dried particles into an oral dosage form.

- 2. (Currently amended) The method of claim 1, further comprising the step of adjusting the pH of said working solution to solubilize said an water-insoluble azole antifungal active agent prior to said providing step.
- 3. (Previously presented) The method of claim 1, wherein said working solution further comprises a surfactant.
- 4. (Previously presented) The method according to claim 1, wherein said single phase working solution has a viscosity of from 10 2000 mPa.s during said combining step.
- 5. (Currently amended) The method according to claim 1, wherein the ratio of an water-insoluble azole antifungal active agent—to water-soluble polymer in said working solution is from 1:0.5 to 1:20 on a weight: weight basis.
- 6. (Previously presented) The method according to claim 1, wherein the ratio of solvent to water in said working solution is from 50:50 to 95:5 on a weight: weight basis.

- 7. (Currently amended) The method according to claim 1, wherein said an water-insoluble azole antifungal active agent comprises active agent in amorphous form.
- 8. (Canceled)
- 9. (Withdrawn) The method of claim 1, wherein said active agent is selected from the group consisting of saquinavir, cyclosporine and paclitaxel.
- 10. (Withdrawn) The method of claim 1, wherein said active agent is saquinavir.
- 11. (Withdrawn) The method of claim 1, wherein said active agent is cyclosporine.
- 12. (Withdrawn) The method of claim 1, wherein said active agent is paclitaxel.
- 13. (Withdrawn) The method of claim 1, wherein said active agent is subject to the proviso that sparingly water soluble antifungal agents are excluded there from.

- 14. (Canceled)
- 15. (Previously presented) The method according to claim 1, wherein said alcohol is selected from the group consisting of methanol, ethanol, propanol, butanol, and mixtures thereof.
- 16. (Previously presented) The method according to claim 1, wherein said water-soluble polymer is selected from the group consisting of hydroxypropyl methylcellulose, methacrylate, hydroxypropylcelluclose, polyvinylpyrrolidones, dextrins and maltodextrins.
- 17. (Currently amended) The method according to claim[[1]]_3, wherein said surfactant is selected from the group consisting of Sodium Lauryl Sulfate; Polysorbate 20, 40, 60, 80; Polyoxyethylene glycolated natural or hydrogenated vegetable oils such as polyoxyethylene glycolated natural or hydrogenated castor oils (Cremophor ®), Poloxamer, Polyoxyethylen 50 Stearate, Propylene Glycol Monostearate, Sorbitan Monopalmitate, and Sorbitan Monostearate.

- 18. (Previously presented) The method according to claim 1, wherein said core particles comprise microcrystalline cellulose spheres.
- 19. (Previously presented) The method according to claim 1, wherein said core particles comprise mannitol spheres.
- 20. (Previously presented) The method according to claim 1, wherein said core particles are from 100 to 1000 micrometers in diameter.
- 21. (Previously presented) The method of claim 1, wherein said working solution is essentially free of methylene chloride, and said oral dosage form is essentially free of methylene chloride.
- 22. (Previously presented) The method of claim 1, wherein said drying step is followed by the step of coating said spheres with an external coating.
- 23. (Previously presented) A pharmaceutically acceptable particle produced by the process of claim 1.

- 24. (Withdrawn) The particle of claim 23, wherein said active agent is selected from the group consisting of saquinavir, cyclosporine and paclitaxel.
- 25. (Withdrawn) The particle of claim 23, wherein said active agent is saquinavir.
- 26. (Withdrawn) The particle of claim 23, wherein said active agent is cyclosporine.
- 27. (Withdrawn) The particle of claim 23, wherein said active agent is paclitaxel.
- 28. (Withdrawn) A pharmaceutically acceptable particle comprising:
 - a central rounded or spherical core comprised of a core material; and a coating film formed on said core, said coating film comprising a water-soluble polymer and active agent; with said particle comprising, by weight, from 5 to 40 percent active agent; from 10 to 80 percent particle core material; and from 10 to 80 percent water-soluble polymer;

and with said particle containing less than 200 ppm methylene chloride.

- 29. (Withdrawn) The particle according to claim 28, wherein said active agent comprises active agent in amorphous form.
- 30. (Withdrawn) The particle according to claim 28, wherein said active agent is selected from the group consisting of protease inhibitors, proton pump inhibitors, oligopeptides, statins, antibiotics, antifungals and antineoplastics.
- 31. (Withdrawn) The particle according to claim 28, wherein said core material comprises microcrystalline cellulose.
- 32. (Withdrawn) The particle according to claim 28, wherein said water soluble polymer is selected from the group consisting of hydroxypropyl methylcellulose, polymethacrylate, hydroxypropylcellulose, polyvinylpyrrolidones, dextrins and maltodextrins.

- 33. (Withdrawn) The particle according to claim 28, wherein said particle further comprises an external coating formed on said coating film.
- 34. (Withdrawn) An active agent oral dosage form comprising a pharmaceutically effective amount of particles according to claim 28.
- 35. (Withdrawn) The dosage form according to claim 34, wherein said dosage form contains from 5 to 500 milligrams of active agent.
- 36. (Withdrawn) The dosage form according to claim 34, wherein said dosage form is a hard-gelatin capsule.
- 37. (Withdrawn) The dosage form according to claim 34, wherein said dosage form is a tablet.
- 38. (Withdrawn) The dosage form according to claim 34, wherein said dosage form is free of lipid or oil solvent.

- 39. (Withdrawn) A method of treating a disorder in a subject in need thereof, comprising orally administering to said subject an oral dosage form according to claim 34 in a pharmaceutically acceptable amount.
- 40. (Withdrawn) The method according to claim 39, wherein said oral dosage form is administered to said subject under fed conditions.
- 41. (Withdrawn) A method according to claim 39, wherein said oral dosage form is administered to said subject under fasted conditions.
- 42. (New) The method according to claim 1, wherein the water-insoluble azole antifungal active agent is ketoconazole.